4 Active packaging in polymer films M.L. ROONEY

4.1 Introduction

Polymers constitute either all or part of most primary packages for foods and beverages and a great deal of research has been devoted to the introduction of active packaging processes into plastics. Plastics are thermoplastic polymers containing additional components such as antioxidants and processing aids. Most forms of active packaging involve an intimute interaction between the food and its package so it is the layer closest to the food that is often chosen to be active. Thus polymer films potentially constitute the position of choice for incorporation of ingredients that are active chemically or physically. These polymer films might be used as chosure wads, lacquers or enamels in cans and as the waterproof layer in liquid cartonboard, or as packages in their own right.

The connection development of active packaging plastics has not occurred evenly across the range of possible applications. Physical processes such as microwave heating by use of susceptor films and the generation of an equilibrium modified atmosphere (EMA) by modification of plastics films have been available for several years. Research continues to be popular in both these areas. Chemical processes such as oxygen scavenging have been adopted more rapidly in sachet form rather than in plastics. Oxygen scavenging sachets were introduced to the Japanese market in 1978 (Abe and Kondoh, 1989) whereas the first oxygen-scavenging beer bottle closures were used in 1989 (see Chapter 8). The development of plastics active packaging systems has been more closely tied to the requirements of particular food types or food processes than has sachet development.

This chapter surveys the range of polymer-based active packaging processes that have been reported, their chemical or physical basis and their application to foods and beverages. Attention is given to opportunities for, and obstacles to, either commercialisation or extension of the current range of application. Some processes which have become economically important are treated individually in other chapters.

Oxygen scavenging

The removal of oxygen from package headspaces and from solution in foods and beverages has long been a target of the food technologist. Introduction of vacuum packaging and inert-gas flushing has provided solutions for some

Table 4.1 Food characteristics influenced by oxygen scavengers

Torgets	Moulds, aerobic bacteria	Insects, larvae, eggs	Rancidity, pigment/numerat loss, twowning	Respiration
Characteristic	Microbiological status	Infestation	Cheancal degradation	Physiological changes

of the problems of distribution of oxygen-sensitive foods as described by Brody (1989). However, the opportunity to improve on the benefits gained by application of those technologies, as well as the chance to treat the problems of distribution of foods individually, has led to the current interest in oxygen-scavenging plastics.

The properties of foods that can be influenced by the presence of oxygen scavengers are shown in Table 4.1. The growth of moulds is particularly important in dairy products such as cheese and in bakery products. Oxygen levels of 0.1% or lower are required to prevent the growth of many moulds. Bacterial growth and the growth of yeasts can be a problem in high water-activity foods including meats and prepared dishes, as well as in juices. Oxygen scavengers can prevent oxidelive damage to flavour and colour in a wide range of foods. Likewise, they can maintain atmospheres with oxygen concentrations too low for insect survival in agricultural and horticultural products. The list in Table 4.2 is indicative of the range of foods which could benefit from oxygen scavenging, their type of packaging and the forms in which the scavenger might be applied. The package converter can decide the radure and quantity of active components used in plastics packaging. This allows the opportunity for tailoring the packaging to the requirements of the product.

Table 4.2 Potential applications of oxygen scavenging plastics

Product	Packaging	Сомровея
Aseptic liquids	Cartenboard, bag-in-box	fitta, costing, adhesive, ink
Bakery products	Flexibles	fibr, etc.
Borrages	Flexibles, bag-in-box	Sirs, etc.
Becr	Crown seal liners	Resin, organosol
Clience	Haibbs	film, etc.
Coffee, rea	Closures, flexibles	fiba, etc., resin
Cereals	Rexibles	fibm, etc.
Dried Fruit	Rexibles	film, ekc.
Dried Fuods/Nuts	Plexibles, closures	film, etc., sesin
Fried snacks	Hexibles	film, etc.
FruitVegetables	Flexibles	film, etc.
Milk powder	Plexibles	film, etc
Meu - frah	Hexibles	film, etc
- baccared	Flaribles	film, etc.
Pastourised liquids	Closures, bottles	resin
Retorted foods	Can lacquees, trays, tidding	regin, film, etc.

which are of increasing importance in consumer societies. Some are suitably packaged using existing processes. However, quality can often be retained longer if residual oxygen is removed. This would allow use of different packaging materials and distribution systems. Some foods, and particularly Wagner (1990) lists a wide range of oxygen-sensitive prepared foods gies in order to allow use of the full range of desired distribution systems. This is particularly important when reduced levels of additives have been beverages, cannot be stabilised adequately with existing packaging technolochosen for regulatory or marketing reasons.

Koros (1990) has set out the maximum quantity of oxygen which a generalised range of foods can take up and still have a shelf-life of one year. These quantities are generally in the 1-200 mg/kg range. Abe and Konooh (1989) have shown the need for oxygen removal by in-pack systems when the economic limit of around 0.5% is reached in the general case. This figure can vary in practice as residue levels of around 2% are often encountered when form-fill-seal (ffs) gas flushing is used commercially. Alternatively, less than 0.1% oxygen can be found in vacuum packs of beef primals where muscule respiration and bacterial action scavenge oxygen.

The most appropriate method of removal of oxygen from a food package depends on the nature of the food, its prior processing and the packaging machinery and the way it is distributed. The factors which may need to be considered, and estimates of efficiency when sachets are used, are summarised in Table 4.3 which is based on a similar table devised by Hirata (1992). who compared sachet technologies with vacuum and gas-slush plastics are to minimise the materials cost by matching the quantity of packaging systems. The expected benefits of use of oxygen-scavenging scavenger to the need, and to keep the filling speed high.

4.2.1 Forms of oxygen-scavenging packaging

to the inner wall of the package. This is done using the Mitsubishi Ageless Sachets merely inserted into the food package constitute most of the present systems in commerce. Alternatively, the scavenger can be hot-melt bonded scuvenger sachet attached to the lid of the steamed rice packages In-pack oxygen scavenging involves use of a variety of forms of scavenger.

Table 4.3 Comparison of headspace oxygen removal systems

	System	Residual O. kPa	Capital	Film	Filling
١.	Vacuum	¢1.5	High	Hisk	nowle 1
م	X, Flesh	1-2	Medium	High	Medium
u	9 + 0	< 1.0 < 1.0	High	High	*
Ð	Scavenger	×0.1	_ 	Medium	High
ا،	P+4	< 0.1	Medium	· Medium	Medium

manufactured in Japan by Ajinimoto under the Take Out brand name. The self-adhesive label concept of US company Multiform Desiceants is used in the Marks & Spencer retail chain's preserved meat packs in the UK.

Beyond the concept of adhesion of sachets to the package lies a wide range of possibilities and it is to this area that much of the recent research and development has been directed. Alternatives include package inserts in the form of cards or sheets, or layers coated onto the inner wall of the package. An example is the beverage crown seal liner currently used in the beer industry. This liner was developed by Advanced Oxygen Technologies Inc. and Zapata Industries Inc. under the name SmartCap and independently by W.R. Grace and Company and under the name Daraforn. Both companies manufacture in the USA. Alternatively an iron-based system has Japan. The industrial development and history of oxygen-scavenging been developed under the name Oxyguard by Toyo Seikan Kaisha Ltd in closures is discussed in detail in Chapter 8.

or dispersed in a packaging plastic or the plastic may be made from a Oxygen-scavenging films or other plastics packaging materials are alternatives to sachets. Low molecular weight ingredients may be dissolved polymeric scavenger. The scavenger may be incorporated into a solid which is dispersed in the plastic or may be introduced into various layers of the package such as in the form of adhesive, printing ink, lacquer or enamel, such as found in cans.

package to the film may isolate the sachet from areas of oxygen entrapment Sachets can be a highly efficient form of oxygon scavenger but their nature does not favour contact with liquid foods or where cling of the or ingress. In such situations it would be preferable to have the scavenger in the packaging material. This allows all exposed surfaces of the food to be deoxygenated or protected from oxygen ingress by permeation.

1.2.2 Plastics packaging as media for oxygen scavenging

cakage. Abe and Kondoh (1989) recommend that Ageless sachets should be day. Such requirements therefore rule out the common heat-seal polyniers Oxygen-scavenging packaging has to date been applied only in packs which have inherently a substantial barrier to oxygen ingress by permeation or used in packages with an oxygen transmission rate less than 20 ml/m3/atm/ and thin layers of the more mediocre barriers like polyester (PET) and nylon 6. However, there can be situations such as in transportation of beef printal cuts in which a shorter period of absolute barrier to oxygen is desirable and these plastics might be used as the barrier. The patent of Speer and Roberts (1993) describes a system involving oxidation of poly(1.2-butadiene) and appears to be directed at that market.

package, as in the case of the PET or PVC bottle, or jar or cup, but are more The commonly found oxygen barriers may be in the form of a single-layer

Table 4.4 Permetablety coefficients P × 10¹⁴ (cm² rum cm⁻² soc⁻² cm⁻¹ Hz) @ 30°C

		of the color in .	
Film material	ž	ď	60
Polyvinylidose chloride	0.0094	0.053	0.29
Polyesiar (Mylar A)	0.05	0.22	1.53
Alyamide (Nylva 6)	0.10	0.38	9:1
Polycatylene ($d = 0.960$)	2.7	901	S
Polychylene (d = 0.922)	<u>e</u>	55	352
Aysiyrene	2.9	Œ.	88
Odyvinykhloride	0.40	1.2	0

(Reproduced from Paine, 1962, with permission.)

likely be part of a multilayer package. Table 4.4 shows the oxygen permeability of a range of polymers. A more extensive tabulation of film perneability values can be found in the review by Bixler (1971)

oxygen scavenger is molecularly dispersed in the polymer it is available to particular advantage of such polymers is that they allow rapid diffusion of permeability films such as plasticised PVC and polyethylene. Where the oxygen in its entirety, unlike there case where solid particles are used. The oxygen and water (at elevated temperatures) from the headspace or food in to the reactive ingredients. Such a characteristic of the polymer helps offset the disadvantage of plastics-film-based compositions in comparison with sachets where the scavenging powder has a large reactive surface area exposed. Polyethylene and polyvinyl chloride (plasticised) are nominated as Oxygen-scavenging compounds can be dispersed or blended in highthe reaction medium in crown closure liners for beverage bottles such as for beer, as described in detail in Chapter 8.

The use of plastics as media in which to disperse or dissolve oxygen scavengers places a severe limitation on the number of reactions which can be involved in the scavenging process. Whereas molecules of the size of oxygen or water can diffuse at an adequate rate, larger molecules behave as if they were immobilised. The fact that a molecule can migrate fast enough to fail food-contact regulatory tests does not indicate that it can be used in an oxygen-scavenging system. Thus the use of iron oxidation in polymers has been a challenge for some years requiring all reagents to be intimately mixed. A further challenge is to establish whether the breakdown of iron particles on oxidation occurs as freely in a polymer matrix as it does in powder form in sachets. Labuza and Breene (1989) note that virtually all of the iron in commercial sachets is available for oxygen removal.

The high gas and vapour permeability of the common heat-seal polymers as those of Hofeldt and White (1989) and Farrell and Tsai (1985). The allows them to be used as reaction media for oxygen scavenging in presence of the oxygen scavenger in the heat-seal layer allows maximum aminates. Indeed this has been proposed in a large number of patents such advantage to be taken of the full thickness of the physical barrier to oxygen xenneation of the barrier layer. However, if the scavenger is incorporated

into the burrier layer as in the Ox-bar bottle of CMB Technologies plc (Rolland, 1990) that portion closer to the outside becomes exposed to relatively large announts of oxygen.

reaction can require the presence of an additional species as with the rusting The permeability of the rolymer medium in which the scavenging reaction occurs need not be the limiting variable determining scavenring rie. Depending upon use chemistry involved, the oxidation reaction can be inherently slow, as with crystalline suffites at room temperature or the of iron. The permeability of the heat-seal layer to both oxygen and water vapour can be limiting as in the mixed solfite/acetate layer in retort ponches patented by Furrell and Tsai (1985). In this case the water is needed in ubstantially larger quantity than the oxygen in order to dissolve the deliquescent potassium acetate in which solution the potassium sulfite dissolves and reacts with oxygen.

woven psyotefins. Several recent patent applications describe claims of enhanced availability to the package atmosphere of granular reagents, as in that of Mitsubishi Gas Chemical Industry Co. Various sulfites can be held in a fibrous layer sandwiched between, for example, one layer of foil, and a second of a plastic or paper with an oxygen permeability greater than 7000 ml/m2/day atm. A second patent describes a package consisting of three layers. The outer layer is a plastics film, the middle is a perforated or nonwoven layer containing an oxygen scavenger and the inner layer of the package is a microporous film which resists the flow of liquids through its porcs. This is claimed to be useful for packaging liquid foods (Ohtsuka et An approach to overcoming any limitation on scavenging rate by the heatseal layer's permeability has been to use microporous polymers such as nonal., 1984). Such films cannot be used where transparency is required but may have application in many forms of packaging.

Traditionally, poly(vinylidene chloride) copolymers have provided a nylons, most oxygen barriers are now water sensitive. The permeability to mers (EVOH) and poly(viny) alcohol) means that. together with traditional exposed to a relative humidity change from about 40-100%. Such a change may render a material which is suitable for packaging an oxygen-sensitive water-insensitive oxygen barrier when used as a layer in laminates, coated films or coextrusions. The introduction of ethylene-vinyl alcohol copolyoxygen or EVOH copolymers increases approximately 10-fold when food at low relative humicaty into one which is most unsuitable at high relative burnidity.

The plasticising effect of water on the barrier properties of EVOM (or other hydrophilic barriers) is time dependent, especially if the hydrophilic layer is protected by a water-barrier layer such as polypropylene as in the case of retortable lunch-cups. When such retortable packs containing a wet food such as an entrée are subjected to steam retorting, water is absorbed by the EVOH in such large quantities that the barrier layer becomes quite

polypropylene layer hecomes wery slow on cooling, so the oxygen (990)) and mica platelets to the EVOH (Bissot, 1990) is used to reduce this permeable to oxygen. The rate of water release through the outer pernicability can remain elevated for many weeks (Tsai and Wachtel, 1990). Although addition of desiccants to the polypropylene (Tsai and Wachtel, impact there is an opportunity here to include an oxygen-scavenger layer in the coextrusion to absorb the oxygen, particularly during the period of enhanced permeability. In fact there has been a recent news report of the introduction of an oxygen scavenger into such packages by Toyo Seikan Kaiska Ltd. Thus active packaging has the potential to contribute to sulving the permeability problem in two ways, providing: desiceants which absorb water in polypropylene; and oxygen scavengers which remove oxygen when it does pass through the hydrated barrier layer.

Whereas elevated temperatures and high humidity have been used to advantage in the research of Farrell and Tsai (1985), the effect of emperature on the performance of oxygen scavengers in polymer-based films has been reported in only rare cases. W.R. Grace (1994) bave investigated the effect of low temperatures on their (optionally photopolymers with glass transition temperatures below -15°C, scavenge at least sensitised) metal-catalysed oxidation of syndiotactic poly(1, 2-butadiene). They have found that this polymer, and certain other low crystallinity 10 mVm²/day O, at 10°C or lower. The photoinitiated system described by Rooney (1994) has been shown to function at 0°C.

4.2.3 Brief history of oxygen-scavenging films

Kuhn et al. (1970) and by Warmbier and Wolf (1976). These systems were based on the earlier work of King (1955) and Abbott et al. (1961) who applied palladium metal to the inside surface of can lids. The cans were sushed with mixtures of hydrogen (8%) in nitrogen to give a mixture in The initial oxygen-scavenging packaging film was a multilayer described by which the residual oxygen could react with hydrogen to form water on the palladium surface.

The reaction of hydrogen with residual oxygen on palladium has been taken further by Hayashi et al. (1986) who vacuum-metallised polyester film When a bag made from this laminate was filled with 500 ml of a mixture of hydrogen-nitrogen 8:92% by volume, this layer was found to be effective in suitable for packaging high-valued items such as probes for an oxygen analyser, although production of a thinner metal layer may change future at 2.5 × 10⁻⁴ moles of Pd/m² and laminated this to high density polyethylene. catalysing the conversion of oxygen to water, reducing the oxygen content from 3-4 to 0.4% in 1 day. Due to the expense of this process it would be

The earliest investigation involving reaction of oxygen in homogeneous

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Holland and Shorter (1981) and by Rooney (1981, 1982). This technique takes advantage of the ease of excitation of oxygen from its ground state to its first singlet excited state and has shown that use of polymers as polymer films was described by Rooney and Holland (1979), Rooney, scavengers or reaction media need not inhibit rapid oxygen removal.

Very little investigation of the chemistry of axygen-scavenging films has been published in peer-reviewed journals. However, numerous patents and some conference proceedings give sufficient detail to allow a comparison of the systems reported. The most evident trend in oxygen-scavenging system patent applications for compositions and designs based on plastics. Very few of these have involved actual polymer exidation but rather have required the reactive ingredients to be dispersed within the polymer matrix or to be sandwiched between film layers. An examination of patent applications worldwide gives the results shown in Figure 4.1 which shows the numbers development during the past 20 years has been the increasing importance of of initial applications for new compositions or designs without consideration of whether additional applications for the same idea have been lodged in other countries. The histogram shows that whereas initially only sachet technologies were considered, there was a slow growth in the number of plastics-based systems devised until the numbers were equal for both types of system in 1993-94.

overall increase in the interest in oxygen-scavenging systems, not from a The increasing number of plastics-based systems results from a substantial decrease in the numbers of sachet technology applications. This has resulted from a more lateral approach to potential reactions coupled with approaches

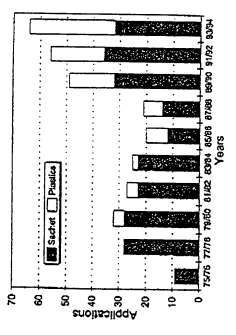


Figure 4.1 Patent applications for oxygen scavengers involving sachets and plastics.

as that of W.R. Grace and Co. for oxidation of squalenc with a transition as oxygen scavengers. The use of bifunctional antioxidants at up to 2% in applications for systems involving oxidation of carbon-carbon double bonds In each case the transition-metal catalysed oxidation results in development of odorous compounds such as low molecular weight aldchydes which are iron oleate, calcium carbonate and active clay are mixed with activated carbon to give a solid oxygen scavenger (Toppan, 1992). The advantage of this system is that water is not needed as a reagent. Some compositions, such metal catalyst (Ebner et al., 1992), appear not to include an adsorbent for odorous products. Some patents involve claims of conventional antioxidants rigid poly(vinyl chloride) was claimed to reduce the permeability of that polymer 20-fold although the period of effectiveness was not reported (Wijbrief, 1971). These antioxidants are normally associated with reactions of primary products of reaction of molecular oxygen with polymers, so the overcoming previous deficiencies. This is particularly evident in patent in small molecules like squalene and fatty acids or in polymers like rubbers. adsorbed by zeolites, carbon or other adsorbents. In one example linseed oil, mode of action of this process is still uncertain.

W. R. Grace and Co. has applied for patents for the use of ascorbic acid liners (Hofeldt and White, 1989). This process relies on the presence of water from the food or beverage as well as the presence of isoascorbic acid and follows the patent of Farrell and Tsai (1985) who patented the between the layers of a retortable pouch structure. Several other applications for oxygen-scavenging plastics containing ascorbic acid have subsequently dispersed into plastics such as the common hear-seal plastics or closure or a metallic sulfite such as sodium sulfite. The process is now commercial sandwiching of either a sulfite alone, or one mixed with potassium acctate, been lodged. One combines oxygen scavenging with the antimicrobial action of a silver zeolite (Shimagawa Nenryo KK, 1992).

blended (Cochran et al., 1991). An additional catalyst in the form of a CMB Foodcan ple has developed a novel system for use in blow-moulded bottles made from PET into which up to 7% MXD-6 nylon had been polymer-soluble cobalt salt such as cobalt stearate was necessary to cause the nylon to react with oxygen.

any form of active packaging, oxygen-scavenging plastics are designed to achieve a specific effect - in this case protection of a packaged food from oxygen. As it turns out, the shelf-life of carbonated beverages like beer is limited by the loss of carbon dioxide by permeation. Thus the effect of the oxygen-scavenging bottle would be to change the nature of the limiting An example of the performance of a bottle made from the Ox-Bar The results suggest that beer held in such a bottle would be saved from oxygen-barrier performance functionally equivalent to that of glass. As with oxygen ingress via the bottle wall for at least 12 months thus providing (Trademark of CMB Technologies plc) has been given by Folland (1990).

variable if used in those countries where beer has a high level of carbonation.

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This process has not reached the market place since the reaction products need to be defined in greater detail. What is particularly interesting in this case is that the use of a cobalt salt catalyst in PET appears to have satisfied iome national regulatory authorities.

4.2.4 Chemistry of oxygen scavenging

be stable in the oxygen-rich environment of air prior to use. This has lew methods of activation appear either in the patent literature or in Unlike most other forms of active packaging, oxygen-scavenging films must presented a problem to chemists formulating such systems and surprisingly commercial practice.

often occurs on the premises of the beverage filler. Thus the catalyst and the oxidisable substrate can be kept apart until the bottle is blown, as in the Ox-Bar process developed by CMB Technologies plc. In some instances activation might not be a necessary consideration if the package is prepared from all constituents immediately before filling. Such packaging systems would include blow-moulding of beverage bottles which

A further factor is the nature of the packaging material carrying the oxygen scavenger. In the case of the PET bottle, containing MXD-6 nylon blended into the PET for instance, the oxygen permeability of the bottle is so low that a delay between blowing and closure is quite reasonable. A general guide to such circumstances is that if the plastic material actually carrying he oxygen scavenger is exposed to the outside air during its life as a package the system can probably be chosen not to have an additional activation step.

The systems which have been developed for use (or prospective use) in coatings, laminations or other plastics layers with high oxygen vermeability re normally activated by one of the following mechanisms.

- Supplying a reagent on package filling, viz. hydrogen or water
 - Supplying water as a solvent or swelling agent on filling.
 - Continuous exposure to light as energy source.
 - Brief exposure to light for activation of o chain reactions, e.g. autoxidation
- o rapid photoreduction of scavenger precursor

can come from the food uself as described in the Oxyguard process of Toyo Scikan Kaisha Ltd, or in the variety of patents referred to in Chapter 1. This In each of the above processes several variants exist. For instance, the water requirement is elaborated in the discussion on sachet technologies in Chapter Water needed as a reaction solveut or to burst micro-capsules can be supplied from the retorting steam in the case of retortable plastics packages which are processed at around 120°C. In this case water vapour permeation can be sufficient from the outside of the pack as well as from the food juseif and lure heat is also a trigger for convinenceneant of the process. Tsai and Wachtel (1990) have shown that EVOH trays can take up as much as 2.8% extra water on retorting, and this diffuses out slowly over a period of some

environments for cultivation of anaerobic microorganisms. Conventionally a 4.2.4.1 Supplying a reagent. The oxidation of hydrogen by molecular oxygen requires metal catalysis at room temperature. Indeed such a process has long been used by microbiologists to generate low-oxygen-content mixture of hydrogen (8%) in nitrogen is either flushed into the cultivation of an air headspace. The surface of palladium metal either as mesh or chamber or hydrogen is generated in a sealed anacrobic jar in the presence deposited on porous alumina catalyses the reaction to form water.

The application of this process to packaging by King (1955) and Abbout et al. (1961) was particularly successful for removing oxygen which desorbs from canned spray-dried milk powder. The hydrogen content of the flushing gas is limited to 7% or less to avoid the risk of explosion. It should be noted that an explosion limit of 6.3% bydrogen is specified in plants for production of chlorine by electrolysis.

The laminate bag described by Warmbier and Wolf (1976) consists of a polyester outer layer bonded to aluminium foil then Surlyn. Between this Surlyn and the heat-seal layer, also of Surlyn, is sandwiched a layer of hydrogen, through the Surlyn to react on the catalyst surface. The quantity of water which was formed was calculated by the authors to be insufficient to powdered alumina upon which palladium metal has been deposited. The bag was flushed with the above-mentioned H2N2 mixture before heat sealing. On storage the milk desorbed oxygen which diffused, together with the marketed as Maraflex 7F by American Can Company, with foods although affect the food. There appears to have been very little use of this system, it was used in the US space programme.

The oxidation of iron in the presence of electrolytes has been well established in the sachet technologies reviewed in Chapter 6. Where pakents describe incorporation of treated from powder into plastics, water ingress is sufficient water on to the iron surface to carry out the reactions described in generally needed. Some specify the need for such materials to be used under retorting conditions. This is probably for the purpose of introducing Chapter 6. It is doubtful whether an adequate rate of oxygen scavenging can be achieved when the polyolefin hear-seal plastics are used as the reaction medium at room temperature.

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Some packaging applications potentially permit the use of scavenging components as heterogeneous additives to the packaging material. Particularly where reagents are in the crystalline form or coated with a low permeability substance, the rate of reaction with atmospheric oxygen can be acceptably low for incorporation into packaging. When the package is exposed to high humidity and elevated temperatures the rate of transmission of water from citlex the food or the outside (e.g. as steam) can be sufficient to dissolve the 4.2.4.2 Supplying water as a solvent or swelling agent. reagent thus enabling oxidation to proceed rapidly.

An early example of the dissolution of a salt was patented by Farrell and Tsai (1985). They incorporated a mixture of potassium acetate and sodium sulfite crystals (or potassium sulfite alone) between the barrier and hear-seal layers of a retort pouch laminate which had five layers overall. The potassium acctate is deliquescent and absorbs sufficient water to dissolve when the food-filled pouch is retorted. The sodium sulfite dissolves in the polassium acetate solution and reacts with oxygen diffusing into this solution from the food and especially from the retort atmosphere. The water vapour permeability of the farninate can increase 1000-fold from 21°C to 121°C. The sulphite reacts with oxygen as in Equation 4.1.

$$2Na_2SO_3 + O_2 \rightarrow 2Na_2SO_4 \tag{4.1}$$

Although this process can result in the presence of an aqueous solution in the laminate, it should be possible to consider use of some form of binder for the solution after retorting. A process such as this which requires both heat and high humidity is really limited to applications which involve recorting or substantial heat-treatment of the packaged food.

complexation agent in separate microcapsules within a polymer film (Zenner A recent patent describes the incorporation of a metallic salt and a et al., 1992). On exposure of the film to an atmosphere of high relative humidity, as in the headspace of some food packs, the microcapsules absorb water and swell. This results in bursting of the microcapsules and mixing of the complexing agent and the salt under moist conditions. The resultant metal chelate forms an oxygen adduct with a binding constant greater than 109M-1. The presence of such a strong adduct is effectively equivalent to oxygen removal by means of an irreversible reaction.

The reaction of oxygen with a metal complex can also involve a weak This type of reaction is the basis of blood oxygenation and can frequently be reversed, at least to some extent (see Chapter 1). This type of oxygen coordinate bond without converting the metal ion to a higher oxidation state. absorber was the subject of extensive research by Aquanauties Inc in the USA, the predecessor of Advanced Oxygen Technologies Inc.

Ascorbic acid and isoascorbic acid react readily with oxygen, more apidly at high pH. The use of metallic salts of these acids is described in

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patents from many sources. The cliemistry of oxidation of ascorbic acid is iterature. It has been observed in the author's laboratory that cellulose acetate films containing dissolved ascorbic acid turn light-brown in colour accepted scavenging compositions have not been described in the scientific complex, consisting of a series of consecutive and concurrent reactions, depending upon the conditions. The steps involved in the commercially after standing in darkness under ambient temperature and humidity conditions for one to two years.

The steps involved in ascorbic acid oxidation in solution have been reviewed by Tannenbaum (1976) who summarised the reactions involved in The initial step is the formation of dehydrouscorbic acid and this step is strongly dependent upon pH. Ascorbic acid can be regenerated by reaction use of sodium sulfite in combination with ascorbic acid (and its isomers) in with mild reducing agents such as metallic sulfites. It is particularly interesting that the patent of Hofeklt and White (1989) describes the optional their oxygen-scavenging closure liner for use with beverages such as been There is thus the potential for use of ascorbic acid as an intermediate in the the oxidation of ascorbic acid. A simpler summary is shown in Figure 4.2.

Figure 4.2 Oxidation of accorbic acid.

whether this desirable regeneration occurs to a very great extent in commercial practice without the formation of an aqueous solution as oxidation of sodium sulfire to the sulfate. Whether this occurs in the ambient Given the heterogeneous nature of the reaction mixture it is doubtful or low-temperature storage of beer has not been described in the literature. described by Parrell and Tsai (1985).

As can be seen from the summary in Figure 4.2 one of the reaction products of oxidation of ascorbic acid to dehydroascorbic acid is hydrogen peroxide or initially the hydroperoxy free radical. Further hydrogen abstraction by this radical forms hydrogen peroxide. The use of sodium sulfite, or another reducing agent, would therefore seem to be desirable if the ultimate re-introduction of an oxidising agent is to be avoided. The role of ascurbic acid as a promoter of browning when oxidised in foods is already known (Tannenbaum, 1976).

Grace and Co. and Zapata Inc oxygen-scaveriging closures in common use The use of ascorbic acid or its isomers or salts is the basis of the W.R. in some beer bottles.

state can be brought about by the process of photosensitisation, which avolves transfer of visible light energy to oxygen via the intermediacy of a lye. A polymer film can be the medium for these photosensitised oxygencavenging reactions. Such a film most contain a photoscarsitising dye and an electron-rich oxidisable compound termed a singlet oxygen acceptor. The oxygen-scavenging process occurs when the blm is illuminated with UV, visible or near infra-red irradiation of appropriate wavelengths. The steps in 1.24.3 Continuous exposure to light as an energy source. Some oxiduion reactions which do not occur when oxygen is in its unexcited (ground) the process are shown below, and these occur within the matrix of the oolymer film.

$$D \longrightarrow^{b\nu} *D \tag{4.2}$$

$$^{*}D \rightarrow ^{3}D$$
 (4.3)

$$^{3}D + O_{2} \rightarrow D + ^{1}O_{2}$$
 (4.4)

$$A + {}^{1}O_{2} \rightarrow AO_{2} \tag{4.5}$$

*D
$$\rightarrow$$
D (4.7)

When a suitable photosensitiser, D, absorbs light (hv) it is excited to a converts to the longer-lived triplet excited state, ³D. (Equation 4.3). In this short-lived higher energy singles state, D. (Equation 4.2) which largely orm the dye can pass the excitation to oxygen by the process of triplet-

from the package headspace (for instance) needs to come very close to the excited, immobile dye molecules during the triplet lifetime of the latter. This is of the order of 10-1000 microseconds. Thus, for such a process to be triplet energy transfer (Equation 4.4). The oxygen diffusing into the polymer useful the palymer matrix (such as an inner layer of a laminate) would need to be very permeable to oxygen.

they are within the distance the singlet oxygen can diffuse before it is Once excited to its singlet state the oxygen can react with any electronrich acceptors, A, present in the polymer matrix (Equation 4.5) provided quenched back to the ground state (Equation 4.6). The process occurs only within the lifetimes of the excited scraitiser (Equation 4.7) and singlet oxygen and so requires continuous illumination. The distance singlet oxygen can diffuse before quenching in a polymer matrix is of the order of 100Å depending on the polymer permeability and other factors (Turro et al.,

This chemistry has been used in the laboratory as an oxygen-scavenging initially but becomes diffusion limited when the oxygen partial pressure reaches low values (Rooney et al., 1981). The light intensity used was 2 x 10^{2} -7 × 10⁵ Lux. It was subsequently shown that the rate of oxygen scavenging reaches a maximum at a concentration of tetraphenylporphine above 10⁻³ M but less than 10⁻² M. When the scavenger film was used as a cellulose (Rooney et al., 1981; Rooney, 1982). The sensitisers include erythrosine or meso-tetraphenylporpline and the acceptors are bis(furity of the polymer film is an important determinant of scavenging rate, with Barrers for cellulose acctate (Bixler, 1971). It was also found that the rate of scavenging of oxygen from a pouch by the film is limited by light intensity roll it was found that the rate was dependent upon the length of film in the the ethyl cellulose being a better matrix for rapid scavenging than cellulose acetate. Ethyl cellulose has an oxygen penneability coefficient at 25°C of 7 process where the polymer matrix is, for instance, cellulose acetate or ethyl Barrers (cm³ (STP) cm/cm⁻² s⁻¹ cm⁻¹ Hg \times 10⁻¹⁰) compared with that of 0.7 furylidene)pentacrythritol and ascorbic acid. It was found that the permeabil roll consistent with increased access of oxygen to the film.

rubber can be photo-oxidised at a rate sufficient to bring about rapid oxygen or ethyl cellulose 0.6 M with respect to PEF and 10-3 M with respect to tetraphenylpunjluine. The rubber scavenged oxygen substantially faster than The singlet oxygen acceptor does not need to be a small molecule dissolved in the polymer. It has been shown the double bond of natural scavenging from the headspace of a package. Figure 4.3 shows the rate of scavenging of oxygen from air, 20 ml, in a 10 cm × 10 cm pouch coated on the PEF in the ethyl cellulose. This was interpreted as being due to the higher concentration of double bond (13.5 equivalents/litte) in the rubber han the concentration of furan rings (1.2 equivalents/litre of film) in the the inside with either natural rubber dyed with terraphenylporphine, 10-1 M.

oxygen (24 Barrers) than that of ethyl cellulose (7 Barrers) is also likely to contribute. It is significant that the reaction occurring in natural rubber occurs on the polymer chain and therefore is not dependent on the free minutes. An unpleasant odour was also generated. The oxidation of the (bifunctional) PEF, although the higher permeability of rubber towards rotation of the acceptor. The photo-oxidation of the tacky natural rubber layer resulted in rapid crosslinking as indicated by the loss of tack within rubber continued on dark storage resulting in formation of a brittle powdery

The potential for use of synthetic rubbers in place of natural rubber was investigated in order to determine whether the nature of the rubber monomers affects the rate of scavenging. The results in Figurc 4.4 show the rate of oxygen scavenging by poly(dimethylbutadiene) (PDMB), cispolybutadiene (PB) and cis-polyisoprene (PI). Based on the reactivity of simple low-molecular weight analogues of these polymers, it would be expected that the more highly methylated rubber poly(dimethylluvadiene) would react more rapidly with singlet oxygen. However, the inverse relationship is observed, presumably due to the low oxygen permeability of PDMB of 2.1 Barrers compared with values of 20 Barrers and 24 Barrers for PB and PI. Thus permeability rather than reactivity can be the important

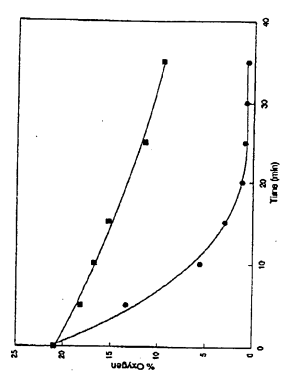


Figure 4.3 Oxygen scavenging by antural nubber e, and PEF 0.5M in eithy cellubase e, both dyed widt barapheaylporphilm, 10°M. Volume of sir, 20 ml, pouch uses 100 cm². (Reprinted from Rooney, 1982b, with permission.)

4.2.4.4 Brief exposure to light for activation.

to activate reactive polymer-based systems has been investigated. The problem, inherent in all oxygen-scavenging systems, of using compounds places an intolerable restriction on the way in which most foods would need to be stored and distributed. It is not surprising therefore that the use of light reactive towards oxygen but stable towards thornal processing of the Continuous exposure of a package to light packaging material needs to be addressed. Chain reactions.

The use of autoxidation of fatty acids for oxygen scavenging has already been claimed but this process seems likely to result in formation of the same Some sachet patent applications describe the use of adsorbents in the sachers volatile oxidation products which are undesirable in foods (Frankel, 1982). to bind volatile off-flavours generated in this way. Mitsubishi Gas and absorbs any odour formed on the oxidation of fatty acids or oils catalysed by transition metal compounds in the presence of alkaline earth bases (Inoue and Komatsu, 1988). Toppan Pritting Company (1992) claimed a similar Chenical Company has claimed that activated alumina, silica gel or charcoal composition involving addition of activated clay. The benefit of such systems is independence from water for the reaction. The research effort put into development of photodegradable plastics packaging over the pust two decades has provided a background to one approach to this problem

Rabek and Ranby (1975) show how a transition-metal metal salt and a photosensitiser dispersed in a plastic can cause it to degrade in darkness ouce exposed to sunlight for some days. This form of photodegradation involves a substantial extent of hydrogen abstraction from the polymer backbone coupled with hydroperoxide formation, particularly on tertiary carbon atoms. Subsequent breakage of the hydroperoxy bond leads to formation of either keto or aldehyde groups which become additional photosensitisers or result in polymer chain scission. It is in the prevention of the latter reaction that the opportunity for development of oxygenscavenging polymers exists.

The scavenging of oxygen including oxidation of hydrocarbon polymers by brief exposure to light of wavelengths less than 750 nm. The optional use (1993). The novelty of their method is that the oxidation process is activated of a photosensitiser is reported to increase the rate of activation, particularly impregnated with salts of transition metals has been described by Speer et al. in the presence of an antioxidard.

As with the photodegradable plastics processes, the chemistry involves laitiation of the free radical process on the polymer chains by changes in the oxidation state of the transition metal ion, preferably cobalt. The redox reaction on the polymer chain involves either hydrogen abstraction in polypropylene or polyethylene or oxygen attack on the double bond in the case of poly(1,2-butadiene).

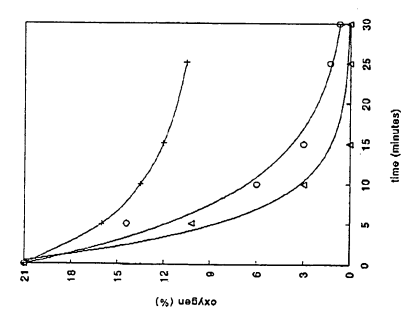


Figure 44 Oxygen scavenging by polydienes, poly(dimethylbutadiene) 1, cis-polyisopene O, cis-polybutediene A. Costings on inner surface of pouches of area 100 cm², volume of air 20 ml.

rariable in an oxygen-scavenging polymer system as was found to he the case with PEF dissolved in cellulose acetate and ethyl cellulose.

The use of multiple dyes in oxygen-scavenging films has been investigated, since illumination with white light of polymer films containing only one dye is wasteful of potentially usable energy (Rooney et al., 1994). It has a natural food colour and a poor photosensitiser (Chignell et al., 1994), was shown to enhance the rate of oxygen scavenging photosensitised by eosine been found that even dyes which are not photosensitisers can harvest energy and transfer this energy to a photosensitiser also in the film. Thus curcumine, n ethyl cellulose. The effectiveness of such a film in suppressing rancidity in sunflower oil has been investigated and found to be substantial at 23% and 37°C (Maloba et al., 1994)

6

activated, the polymer needs to be stabilised against premature initiation in the extruder or moulding machine in package fabrication. Antioxidants this protection and to relard the autoxidation chain reaction for a selected period of time between activation and package filling. Removal of antioxidant from the polymer appears likely to increase the likelihood of aint formation due to unrestricted polymer oxidation. Attention should probably be given to inclusion of odour absorbents in the composition as retains its physical properties even when most or all of its oxidisable groups have been reacted. Since this is a typical autoxidation that is also thermally provide this protection. The quantity of antioxidant is chosen both to provide peroxides or carboxylic acids, in some cases with chain scission. In the case containing the double bond. Speer et al. (1993) report that this polymer The hydroperoxide formed by reaction of either the polymer free radical or by direct attack on the polymer decomposes to form ketones, ableliydes. of poly(1,2-butediene) it is the aim to localise the oxidation to the side chain proposed by Toppan (1992).

The manner in which such a system could be used is particularly broad but one suggested is to laminate or coextrude the reactive layer with an oxygen-barrier layer. Such an application would be appropriate for many packaging processes such as in the storage of cheese, nuts and meats such as beef primal cuts.

The use of such compositions at the low temperatures required for storage of meats and cheese requires that the reactive layer be readily permeable to oxygen. Specr and Roberts (1993) have specified that the compositions containing their oxidisable components should be largely amorphous and have a glass transition temperature below ~15°C.

4.2.4.1. Rapid photoreduction of a scavenger precursor. Research into systems involving this form of triggering indicate that this approach overcomes some of the limitations of other methods already proposed (Rooney, unpublished results).

4.2.5 Chemical barrier to oxygen permeation

Implicit in the use of oxygen seavenging inserts (sachets) or closures is the effect these have on the consumption of oxygen as it enters the product during its storage life. The results in Table 4.5 show the calculated impact of

Table 4.5 Calculated life of chemical oxygen burriers

	OTR	Barrier life
Film	cm2 m-2 day-1 atm-1	(days)
1 005 34 mm	000\$	۲,
Mary Ac 11 M CABC 25 11 mg	£9	×
rei Dimilire Dimil	; 0	362
PVIXC-coated CPP 25 µm	` ;	350
Meislined PET 12 pro/LDPE 25 pro	20	(67)

the oxygen transmission rate of packaging materials on the period of effective oxygen scavenging by a commercial oxygen-scavenging sachet with an absorption capacity of 50 ml of oxygen. The packages are 10 cm × 20 cm and initially contain 100 ml air. LDPE is low-density polyethylene, OPP is oriented polypropylene, PET is polyester and PVDC is poly-(vinylidene chloride). However, in a food-packaging situation where the food is tightly packed, the sachet can be expected to deoxygenate only a small portion of the headspace and combat permeation of oxygen that is accessible to it.

Close-fitting packages such as vacuum packs for block cheese and meats or aseptic cartons of beverages are examples where the headspace is very small and oxygen permeation is the prime cause of quality loss. It is in such circumstances that oxygen-seavenging plastics films are particularly needed. The use of the oxygen-seavenging reaction to intercept oxygen diffusing through the package wall is an example of a chemical barrier as distinct from the physical barrier normally provided by aluminium foil and vacuum metallising or crystalline polynners such as poly(vinylidene chloride).

film of intermediate performance coupled with a chemical barrier. This period of oxygen transmission at a rate less than that found in darkness. In combination was investigated by Rooney and Holland (1979) and the results in Figure 4.5 demonstrate a period of total barrier to oxygen permeation from air at 25°C. A laminate of polyethylene/nylon 6/cellulose acetate containing oxygen-scavenging reagents was used to separate the two cent light resulting in a barrier life of around 30 days was followed by a The use of a chemical oxygen barrier offers the opportunity to cheapen barrier packaging for relatively short shelf-life products such as wholesale or export units or for some fresh foods. This can be achieved by using a barrier compartments of a glass permeability cell. When the cthyl cellulose layer contained 0.5 M bis-furfurylidenepentaerythritol, illumination with fluoresthis case the chemistry used was a singlet oxygen reaction which requires constant illumination to photo-excite the oxygen. Other chemistries can be used to achieve similar reactions in darkness and Speer et al. (1993) identified this application for their photoinitiated chain reactions in rubbers, and Cochran et al. (1990) devised the Ox-Bar process largely for this

The use of oxygen-scavenging sachets to allow the use of a cheaper packaging material has been described by Alarcon and Hotchkiss (1993) and their work is reviewed in Chapter 6.

Although use of multilayers of plastics films appears the most appropriate approach to solving permeability problems, there have been early attempts to carry out scavenging reactions in liquid layers. Oxygen scavenging in liquid layers was one of the earliest processes described in the patent literature. Cook (1969) claimed that bilayers of plastics films separated by a layer of a solution of one of several antioxidants in high boiling point solvents

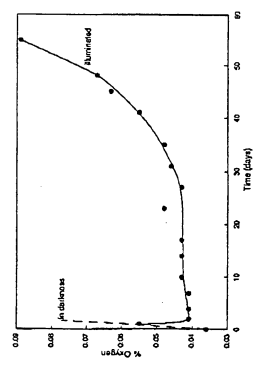


Figure 4.5 Barrier to oxygen permanton of polychylese/Nyton 6/cethulote acrine laminate. Cellulote accuse 26 pan thick containing PEF (0.5 m) and erythrosine, 5 x 10⁻¹ m. Oxygen concentration measured in a cell on the cellulose accuste side of the laminate, air on the other

demonstrated a reduced oxygen permeability. Whereas the mechanism was unknown, all of the common antioxidant classes were claimed to be Use of an aqueous solution of an oxygen-scavenging reducing sulfite as a layer between two polymeric layers has been claimed by Scholle (1976). As oxygen permeability of the laminate. In the case of Cook's system the role appears to have been to improve shrinkable packaging, as for meat, whereas with the organic system of Cook (1969), the purpose was to reduce the the Scholle system was particularly suitable for bag-in-box liners since the packages consist of, at least, a collapsible duplex which is sealed together only at the edges where the liner forms a bag (and around the tap fitting). Such a system requires an oxygen scavenger for oxygen-sensitive products due to the gas space between the two film layers.

43 Moisture control films

Moisture affects the gas and vapour permeability of hydrophilic plastics packaging films (Davis, 1966). Fruit cakes packed in films with an appropriate water vapour permeability can have long shelf-lives because the surface dries somewhat. This creates conditions unfavourable to mould growth. Nitrocellulose-coated regenerated cellulose has been used because it

PULYMER FILM

prevents contamination with moulds and allows substantial water vapour permeation.

There are, however, several sets of circumstances in food packaging where greater selectivity in control of liquid and gaseous water is required. These include:

- Transpiration of horticultural produce.
- Melting of ice, e.g. in fish transportation.
- Temperature fluctuation in high erh food packs.
- Drip of tissue stud from cut meats and produce.

build-up of liquid water. Alternatively, the natural water level in the tissue The problems which arise from these circumstances are the result of can be undesitably high near the surface for microbiological stability under the chosen storage conditions. The results of the presence of water in unwanted quantities include bacterial (yeast) and mould growth as well as fogging of films and mobile blood and tissue fluid. Moisture migration via the vapour place can result in transient formation of regions without adequate preservative or where the a, is high even though the food is packed at lower a.. Some of the ways of addressing these problems via packaging are considered in detail in terms of either liquid water control or humidity

4.3.1 Liquid water control

non-polar chain in the plastic and the polar end group at the interface. The Temperature cycling of high exh foods has led most film manufacturers to use heat-scal plastics with anti-fog additive. These additives are blended with the resin before extrusion and migrate to the surface after film formation. The additives are chosen for their amphiphilic nature with the result is a lowering of the interfacial tension between water condensate and the plastic film. The fog droplets therefore coalesce and form a transparent film on the plastic. This film may even flow on sloped surfaces and gather at the bottom of the pack in extreme circumstances.

Anti-fog treatments are a cosmetic form of active packaging, assisting the customer to see the packaged food clearly. There is no change in the availability of liquid water in the package and this has the potential to cause Other active packaging systems go further in removing liquid water from produce spoilage unless managed by one of the processes described below. food contact.

Several companies manufacture drip-absorbent sheets, which may have a variety of other names. Basically they consist of two layers of a microporous a non-woven polymer, such as polyethylene or polypropylene, between which is placed a superabsorbent polymer in the form of free-flowing granules. The duplex sheet is scaled at the edges and is normally quilted to allow the water absorbent to be held in place rather than aggregating towards one edge of the sheet when tilted towards one edge.

the white foam tray. Given the interest in reducing the volume of solid waste by decreasing foam polystyrene usage, the use of drip-absorbers may become even more necessary to prevent water damage to the more Such sheets are then used either as pads under whole chickens or chicken pieces to absorb drip, thereby preventing discoloration of either the meat or hydrophilic afternatives.

absorbing at least 100 times and possibly as much as 500 times its own A recent patent application in Japan describes a water-absorbent sheet which Large sheets are used for absorption of melied ice in the packaging of seafood for air transportation. The superabsorbent polymer used in the Thermanic@ Pty Ltd sheet manufactured in Australia is capable of 1994). Another product of this type is Toppan Sheet manufactured in Japan. weight of liquid water depending dramatically upon salinity (Malouthi et al., has a high barrier to oxygen for use with fresh meat (Showa Denko KK, 1990). This would necessitate use of a scaled outer pack.

although graft copolymers of starch can also be used. Such copolymers consist of a polysaccharide backbone with synthetic polymer chains saccharide chains is disrupted allowing the starch to exercise its affinity for water by hydrogen bonding. Such polymers tend to become slimy when The preferred polyniers used to absorb the water are polyacrylate salts, radiating from the starch backbone. The strong association of the polyswollen with large amounts of water.

The swelling of the polymer on hydration results in substantial distortion of the duplex sheet, an effect that is controlled somewhat by the quilted seal pattern. This effect can be controlled by choice of the amount and the capacity of the superabsorbent polymer.

The significance of such active packaging in seafood shipment by air lies in the removal of the potential for spillage of salt water from cartons in aircrast holds. The aluminium construction of aircrast makes them easily susceptible to corrosion damage, at great cost.

4.3.2 Humidiny buffering.

allows the food packer or even the householder to reduce the surface a plastic film which is highly permeable to water vapour. An example of this type of product is Pichit manufactured by Showa Denko in Japon. The film duplex is described as containing an alcohol, described by Labuza (1989) as packages is to intercept the moisture in the vapour phase. This approach This can be done by placing one or more humectants between two layers of An entirely different approach to the control of excess moisture in food concentration of water in a food by reducing the in-pack relative humidity. propylene glycol and a carbohydrate, both of which are humectants.

POLYMER FILM

Table 4.6 Water vapour optake by Pichit and Toppan sheet

These	Weight loss fi	Weight loss from petri dish (%)
(b)	Pichit	Toppan skeer
2	1.2	1
~.	2.4	t 0
4	3.6	1.2
స	15.2	~
22	22.0	80

The effectiveness of such a combination material in comparison with a Distilled water 5 g was placed in each of two petri dishes (with lids) and these were enclosed with either Pichit or Toppan Sheet (450 cm²) in a highpartier package. The packages were opened at intervals and the petri dishes were weighed to give the per cent loss of weight of water, shown in Table 4.6. Thus, whereas the superabsorbent polymer has a far greater capacity for liquid water uptake, the humectant based film more rapidly absorbs water superabsorbent polymer absorbent for liquid water is shown in Table 4.6.

pieces of flesh food such as fish or chicken to reduce the a, proximate to the sufficient to allow some of the latter to diffuse to the food surface. If this is the case, some antimicrobial action might be expected. Propylene glycol is Pichit is marketed for home use in roll or single sheet form for wrapping ood. At present there is a lack of experimental verification of the significance of this effect. Louis and de Leiris (1991) suggest two to three days of fresh storage is possible. It has been suggested by Labuza (1989) that the permeability of poly(vinyl alcohol) to propylene glycol may be a GRAS substance in the USA.

Louis (1991) reports the availability of an additional moisture control film which plays the same role as Pichit in the Japanese market. The potential for use of such materials for quality retention in the thome appears to be considerable and this type of film should be subjected to detailed objective evaluation for use in the domestic situation where excess portions of flesh loods need refrigerated storage for some days. The potential for washing the surfaces of sheets like Pichit, followed by re-use, is an attractive incentive for active packaging use in the home provided absence of microbial contamination can be demonstrated.

for use in the distribution of hartlcultural produce. Currently the packages are wholesale fibreboard cartoirs of produce, usually with a polycthylene sbreboard cartons to allow moist produce to be placed directly into the A different approach to humidity buffering has been under investigation recent development has been the water-barrier coating of the inside of carton. Besides the introduction of liquid water with the produce, packing liner or consisting of the very expensive waxed fibreboard without a liner. A nto closed spaces allows the build-up of water vapour. The produce

continues to lose water by evaporation during distribution and the relative humidity in the lined carton can reach close to 100%. Since temperature cycling is very difficult to avoid during handling, there is every likelihood of condensation, and, with this, the growth of microorganisms on fruits and

Two widely different approaches have been taken to buffering the humidity in the cartons in order to prevent condensation while not concurrently causing desiccation of the produce. One is to include microporous bags or pads of inorganic salts and the other is to line the carton with a protected Layer of a solid polymeric humectant.

The approach of using microporous sachets of inorganic salts has been used in the US tomato market in recent years. This was proposed by Shirazi and Cameron (1992) who showed that an equilibrium relative humidity of Hudson (1991). Both these studies involved use of heat-sealed liners which around 95% above produce can be reduced to around 80% by use of sachets of desiceant sult such as sodium chloride. The application was extended by were concurrently being evaluated for their performance to maintain equilibrium modified atmospheres (EMA) generated by the fruit (see Chapter 3). The control of moisture is one of the main impediments to the introduction of EMA packaging. Indeed it is common to package potatoes in perforated sacks to prevent unwanted build-up of condensation.

The most recent alternative involves the use of the carton as the active EMA generation less readily as the humidity is buffered at the interface with the fibreboard. The designs of Patterson and Joyce (1993) involve; an package rather than an insert. This approach lends itself to combination with integral water vapour barrier layer on the inner surface of the fibreboard; a poper-like material bonded to the barrier and which acts as a wick; and a layer highly permeable to water vapour but unwettable next to the fruit or vegetable. The latter layer is spot welded to the layer undemeath. Accordingly, the multilayer of material on the inside of the carton is able to take up water in the vapour state when the temperature drops and the RH nses. When the temperature rises the multilayer releases water vapour back into the carton in response to a lowering of the RII. The condensation control system therefore acts as an internal water buffer. The critical characteristic of the system is the capacity of the wick layer for water.

The performance of this system is demonstrated in Table 4.7 taken from Patterson et al. (1993) which shows the results of a comparison of the free

Table 4.7 Free water in carrot packs on cooling

Puctaging	Cooled from	Cooled from
Palivitulene line	10 to 3°C	22 to 3°C
Condensation aminal comm	10.1 (0.6)*	229 10 20
	0.0 (0.0)*	0.2 (0.2)
Junior error of the meas (1 2 5)		

water found in cartons of carrots with a conventional liner with the cooled to 3°C from either 10°C or 22°C and after 3 days the amount of free condensation control multilayer. The cartons containing carrots, 6 kg, were water in the loxes and on the carrots was measured. The results demonstrate the capacity of such a system for water uptake under circumstances likely to

Shrink wrapping is an alternative approach to the use of active packaging systems for control of condensation on spherical fruits such as citrus. Ben-Yehoshua (1989) has reviewed this field, to which he has made such a large

44 Removal of taints and food constituents

able commercial interest. Hirose et al. (1989) demonstrated the impact of the The interaction of packaging plastics with food aroma has long been recognised, especially through the flavour 'scalping' which is of considernature of the metal ion in Surlyn film layers in aseptic brick-packs on the scalping of limonene from orange juice. In periods as short as 2 weeks at 24°C, almost 30% of the limonene was found in the Surlyn-1601 and 20% in the polyethylene layer in brick packs. Commercially, plastics packaging has not been used to remove selectively components of the flavour or arona of foods which are considered undesirable, but a potential opportunity has been available for over a decade.

Some varieties of orange, such as the Navel, contain a tetraterpenoid of the formula C26H30O3 which is initially present in the albedo but which is extracted into the juice on standing or heating. Thus the juice of such oranges becomes bitter on pasteurisation when this compound, limonin, Processes have been developed for debittering such juices by passing them reaches concentrations exceeding 8-12 mg/kg (Chandler et al., 1968). through columns packed with cellulose triacetate or nylon beads (Chandler

A simple active packaging process was suggested by the same authors, who proposed that since the juice extracts the bitter principle on standing for 24 h, inclusion of the absorbent in the packaging might remove it as it is extracted. To this end they proposed using their absorbents in film form such as cellulose triacetate or as acctylated paper. They showed that a 1-litre plastic bottle coated internally with cellulose acetalo-butyrate reduced the limonin content of 500 ml of juice from 42 to 11 mg/kg after 3 days' refrigeration. Similarly, when cans lined with acctylated filter paper containing juice with 14.9 mg/kg limomin were spin cooked and allowed to stand, the juice was only slightly bitter affer 4-13 days.

It appears that this process has not been taken up commercially although it offers considerable potential for freshly squeezed Navel juice marketing

(Johnson, R.L., private communication). On the other hand, the sorption of imonene oil from packaged juices by heat-seal layers has been the subject of several studies (Mannheim et al., 1989, Hirose et al., 1989). Thus opportunities for active packaging can be closely related to problems of food-package interactions.

A closely related goal had long been achievable commercially in the of sulfur compounds from the food. These sulfur compounds cause the compounds reacting with them before they can diffuse to the tinplate tinplate canning of foods in which protein degradation resulted in the release phenomenon of 'sulfur staining' on the tinplate and so it has been the practice to disperse zinc oxide in tinplate lacquers to intercept such

The remaining methods described in the literature to date for removal of a specific interaction or reaction with a functional group known to be present in the taint or undesirable food component. Two types of taints amenable to taints or off-flavours have largely involved incorporation of ingredients with removal by active packaging have been identified by researchers responsible for current commercial products. These are amines resulting from protein breakdown in fish muscle and aldehydes formed from the breakdown of The formation of aldehydes can make a wide variety of oil-containing foods organoleptically unacceptable well before there is significant damage to the peroxides which result from the initial stages of autoxidation of fats and oils. nutritional or functional properties of the food. Examples of such products would be fried snackfoods such as potato crisps, biscuits and cercal products. Early developments occurred in Japan where there was seen to be need to remove arnine smells from fish which was stored in domestic refrigerators. The amines formed in fish muscle degradation include strongly basic compounds and thus are potentially strong in their interaction with involved incorporation of such acids in heat-seal polymers such as acidic compounds such as citric or other food acids. Hence the carlicst work polyethylene and extruding them as layers in packaging (Hoshino and Osanai, 1986).

ANICO Company Ltd in Japan under the trade name ANICO BAG. Bags A later approach to removal of amines odours has been provided by the made from film containing ferrous salt and an organic acid such as citric or ascorbic acid are claimed to oxidise the amine or other oxidisable compound as it is absorbed by the polymer.

If these materials can be shown to be effective there is an opportunity to Questions which would need to be answered are What is the nature of the products of such reaction and What is their fare. Questions such as these are determine which variables optimise their rate and extent of reaction. of particular interest to regulatory authorities, and the potential for several active packaging systems to generate mobile reaction products is considered

Removal of aldehydes such as hexanal and heptanal from package recently introduced tie layer Bynel IXP101 which is a high-density polychylene (HDPE) resin masterbatch. This masterbatch is blended (2.5-12%) with unmodified HDPE or other linear polyethylenes to form an intermediate layer in coextrusions. It is specified that the heat-seal layer should not be a 'good to excellent gas barrier' (Dupont, 1993). It is interesting to note that the use of a form of active packaging can place headspaces is claimed by Dupont Polymers, Packaging Division, for their restrictions on other components of the packaging. The restriction in this 220°C to avoid fuming. The chemistry of the process is not described but case, and many others, is that the extrusion temperature should not exceed such a process would require the reaction with the aldehyde to be effectively ineversible at least over the temperature range the package is likely to encounter. One such reaction would be the formation of a Schiff base by reaction of the aldehyde with an amino group. The amino group would need to be rather stable to heat and oxygen in order to remain unaffected after the extrusion at temperatures up to 220°C in an air atmosphere.

There may well be a wide range of food constituents which can be components or by chemical reaction with them. A fertile research field would seem to be open especially with liquid foods since solubility and removed by making use of specific interactions with selected packaging diffusion of food constituents in the packaging can be utilised so that the removal process is not limited to compounds with a significant vapour pressure at distribution temperatures. It will be necessary for industry and regulators to ensure that such processes are not used to conceal the marketing of sub-standard or even dangerous products if for instance microbial odours were to be scavenged.

The tainting of foods by compounds originating either in the packaging material itself (c.g. monomers) or outside but permeating the packaging material continues to be a source of problems for the food industry. The approaches have been described in the patent literature. These are formation approaches described above may contribute to their solution but additional of chemical barriers as distinct from physical barriers like aluminium foil or crystalline polymers.

acrylonitrile-butadiene-styrene copolymers or blends being extruded (Tokas, 1979). Addition of myreene is claimed to introduce no new taint Myrcene (7-methyl-3-methylene-1,6-octadiene) has been found to react with traces of styrene or acrylonitrile when the latter are present in while reducing residual levels of the monomers. In another patent, foods packaged in plastics consisting of more than one layer are claimed to be protected against taints from the outside by inclusion of the appropriate adsorbent in the outer layer (Kiru Kogyo KK, 1994). The adsorbent is chosen for expected taints and is kneaded onto the outer layer, apparently retaining the taints there. The period of this type of equilibrium accorption

needs to be established, especially when a package is subjected to to be stored or shipped together with odorous products which are lentestature changes. This may have application in areas where products are inadequately packaged.

A compound which while not a taint is often found undesirable in The potential benefits resulting from such removal are particularly great and packages of respiring horticulural produce is the gaseous auxin ethylone. are discussed separately in Chapter 2.

4.5 Ingredient release

Active packaging materials considered so far have exerted their action on the packaged food by removing unwanted components of either the food or of the headspace enclosed with the food. Another form of interaction is by release of desirable ingredients into the food from the packaging materials or Most are used for their antimicrobial activity, although sulfur dioxide also serves as a chemical stabiliser of colour and flavour by preventing progress of the Maillard reaction which causes non-enzymic browning of products also known as B-thujapricin (Hirata, 1992), derived from cypress bark, is an from inserts packaged with the food. Some substances released commercially or which have been the subject of investigation are listed in Table 4.8. such as dried fruit and wines (Davis, 1975; Davis et al., 1978). Himbkitinl, additional antimicrobial compound specific to the Japan market.

Processing of foods often results in loss of flavour by degradation or evaporation. Another mechanism of flavour loss is the scalping of some tion occurs after packaging. There is, however, the question of whether a flavour compouents by plastics used in packaging (Mannheim et al., 1989). There is therefore the opportunity to replace these lost food constituents by diffusion from the packaging, especially where scalping or flavour degradafood is being sold as fresh when this is not so. This is more a legal matter of consumer protection than a technical one as the question arises of when

Table 4.8 Substances emitted by Active Packaging

Substance	Purpase	Source	Reference
Carbon diuxide	antimicrobial	film	Rooney, 1989
		sucher	Abe, 1990
Ethano!	antimicrobial	sachet	Abe, 1990
Silver ion	nationicrobial	Elg.	Hirata, 1992
Organic acids	antimicrobial	ela ela	Hotchkiss 1993
Suffur dioxide	antimicrobial	sachet	ICI Australia
Benomyl	antimicrobia	E.	Halck and Gary, 1988
Flavours	fortification	elis	Ventor, 1986
finokiris	antimicrobial	2	Abc. 1990
31.1	antioxidant	Sign .	Han et al., 1987
Enzymes	various	E	Buday, 1990

POLYMER FILM

savour addition to a food sabricated from many ingredients becomes confrary to consumer interests.

release flavour precursors in the packaging material. The manufacture of slavour concentrates in the common commodity plastics has been described by Venator (1986). Master batches of plastics with concentrations of up to 40% of the flavour have been marketed with a view to obviate the effects of Release of very few slavours has been investigated to date, although the ease of oxidation of many flavours suggests an opportunity to provide slow-Bavour scalping.

4.5.1 Antiaxidant release from plastics

Two factors acting concurrently are likely to influence the use of packaging nesterials as sources of antioxidants in some foods. The first of these is the need of the industry to respond to pressure by some consumer advocates for reduced use of food additives (Smith, 1993). The second is the renewed food antioxidants in polymer stabilisation replacing some of those developed specifically for plastics. Difaurylthiodipropionate and its base acid, thiodipropionic acid, are approved food additives in some countries and are used interest of plastics resin manufacturers in using natural, or other approved as stabilisers in food-grade polyethylene (Anon, 1992).

evaporation into the package of out flakes. It was found that at 39°C only 55% of the original BHT remained in the film after 1 week. The loss by The potential for evaporative migration of antioxidants into foods from who developed a theoretical model. This work has been taken further by Han et al. (1987) who determined the effect of temperature on both the diffusion coefficient of buylated hydroxytoluene (BHT) in HDPE and the rate of its their packaging plastics has been studied by Calvert and Billingham (1979) After 6 weeks the HDPE film was free of untoxidant and 19% of that originally in the film remained in the cereal. The workers compared the impact of two starting levels in the film on oxidation of the cereal oil and found that with an initial 0.32% BHT there was less oxidation than with outward migration was 70%, and 25% of the BHT was found in the cereal

These results of Han et al. (1987) demonstrate the potential for release of intioxidant into foods provided the rate of diffusion can be matched to the tood's needs. The outward loss can be controlled by use of a layer of film with low permeability to the antioxidant or by use in a closure. In the case of liquid foods or solids with close-fitting packaging the process could be based on diffusion alone and not require the antioxidant to be able to evaporate,

Commercial use of this approach to antioxidant release has been reported by Labuza and Breene (1989) who noted that waxed paper has sometimes

been used as a reservoir for antioxidant release by the US cereal

Other antioxidants might also be used in this way. Lignert and Eriksson (1980) found that Maillard reaction products have a strong antioxidan unction. This work has been extended by Anese et al. (1993). The potential for applying such antioxidants to foods via packaging materials has been suggested (Eriksson, private communication)

perform as oxygen scavengers when used in couting or lamination of The patent of Goyo Shiko KK (1993) describes application of amino acids and saccharides which produce reducing sugars on decomposition and packaging films. These compositions would be expected to undergo the Maillard reactions but their rate would depend upon the thermal conditions used in preparing the packaging and any thermal processing of the packaged product. The inventors nominate liquid foods in cans as a target product area and, given the hydrophilic nature of the polymers involved, it appears likely that Maillard reaction products could be extracted into the food to act as antioxidants as well as scavengers.

viamin E cause less noticeable flavour in distilled water than bottles stabilised with either BHT or Irganox (a bindered amine antioxidant for polymers). There is an opportunity to investigate whether oils can be stabilised by diffusive addition of antioxidant at sustained low levels from It has recently been shown that polyethylene bottles stabilised with the packaging.

The observation of Han et al. (1987) that BHT was lost outwards from HDPE film packs points to an otherwise unrelated opportunity for active other agricultural products for reduced levels of pesticide residues in these products at the time of delivery. Packages such as sacks and lined cartons packaging. There has been constant pressure from importers of grains and containing these products are often attacked by insects during warehousing and transport. The inclusion of low-toxicity furnigants such as pyrethrins in an outer layer of packaging material offers the opportunity to achieve sustained insecticidal activity without substantial addition of fumigant to the food. Highland and Cline (1986) found that polypropylene containing 203 mg m-2 of permethrin provided rather similar resistance to attack by burrowing insects to that provided by polyester which has a harder, slippery surface. Their work involved exposing: polyethykne; polypropylene/polypropylenc/polyethylene laminate, the latter containing permethrin in the outer layer; and polyester/polyethykne pouches of many foods, to four insect populations. The permethrin-treated pouches were resistant to two insect species for 24 months. The effect was not entirely consistent since the treated pouches failed at 24 months with one insect species and were better than polyester with one of the remaining species and worse than polyester with the other. The authors concluded that the order of descending resistance of films to the insects was permethrin-treated film, polyester film, polyethyl-

ene film and worst, polypropylene film. This work nught be developed to provide a low-cost answer to some of the major problems of fumigation, including the cost of repeated fumigations with methyl bromide approximately bi-monthly, as well as reducing the exposure of staff to furnigant application conditions. The pennethrin or alternative treatments would need to be submitted for regulatory approval.

It has been suggested recently that enzymes might be released into foods action. Labuza and Breene (1989) have reviewed the potential for release of bound enzymes into foods. Enzyme inhibitors might also be bound to a film from packaging materials, probably to achieve effects such as antimicrobial surface. An example would be the binding of the inhibitor of methyl esterase to the package surface to bind methyl esterase. The result might be prevention of cloud drop in fresh orange juice (R.L. Johnson, private communication). The subject of enzymes in active packaging is discussed in Chapter 7. The use of enzymes in edible coatings is discussed in Chapter

4.6 Permeability modification

There are a number of circumstances in which it is desirable for the The importance of predicting the requirements of plastic film's permeability been discussed in Chapter 3. It is possible to modify the permeability of a to carbon dioxide and oxygen in packaging of fresh horticultural produce has A variety of semipermeable patches were initially developed by the Hercules company in the USA. Subsequently the effect of sorbed water on cellulosic packaging material to be more permeable to one substance than to another. window in a package and to control gas exchange through this limited area. patches has been claimed to give selectivity in gas permeability matched to the respiration rates of some produce.

There are other substances which need to be selected for entry into plastics packaging. Of particular interest is the ability to selectively transmit ingh humidity and highly impermeable to oxygen at room temperature under snoke flavours through films which are useful as skins for ham and other preserved meats. These are films of polyamide alloy which are highly permeable to water and oxygen under conditions of high temperature and dry conditions (Nishini and Yoshii, 1988). Hirata (1992) described one such 60% RH and a water vapour transmission rate of (WVTR) 60 g/m³/day at type of film as having an oxygen transmission rate of 8 ml/m³/day at 20°C. 40°C, 90% RH. The low oxygen permeability is accessary for colour retertion of preserved meats. This high WVIR should favour rapid transmission of polar flavours. Kureha Chemical Co. (1986) described a polyanide film for this purpose with an OTR in the range 50-300 ml/ml/day for film thicknesses of 5-50 µm. KALYMER FILM

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Current use commercially

ment of the most soundly besed technologies is commercialised. Ethylone scavenging, condensation control and equilibrium modified atmosphere scientific research. This research into equilibrium modified atmosphere packaging has been based on the work of Kader (1980) and earlier workers including Jurin and Karel (1963). Since some of the active packaging technologies currently commercial are scarcely improvements on existing technologies there will probably be significant realignment in the market place. The need for active packaging solutions to problems in the storage and distribution of horticultural produce may well ensure that the developforms of enhanced permeability films have been commercialised for both trade and home use. This success, in spite of a lack of soundly based evidence of effectiveness in many cases, owes much to substantial prior The commercial development of plastics-based active packaging has not application. The major field to date less been borticulture in which several occurred evenly either geographically or in terms of their field of packaging will continue to be emphasised.

processed food field that commercial development of plastics-based systems can be expected to be substantial over the next few years. Perhaps Toyo the semi-aseptically packaged boiled-rice market by Sato Food Industry Co. causing, as is planned, a 100 000 meals/day operation to be more cost Active packaging for processed foods is still based largely on sachet echnologies, with the exception of moisture control packaging. It is in the Scikan Kuisha's planned manufacture of an oxygen-scavenging laminate for Ltd is the first example. This tray has been described as 'epoch making', effective (Anon., 1994).

The substantial impact on the marketplace of oxygen-scavenging erown seals for beverages, metallised polyester microwave susceptors and timetemperature indicator strips is discussed in other chapters.

Regulatory and environmental impacts

regulatory considerations appear to have caused the delay in the introduction Since plastics-based active packaging involves not only changing current regulatory authorities must become involved in many developments. Indeed materials sontewhat but also inclusion of reactive compowents in some cases, of a chemical barrier to oxygen into PET beverage bottles.

The effect of environmental considerations on plastics-based active Commodity films used for produce packaging may become the object of recycling schemes, especially in the European Union, and so additional ingredients will need to be evaluated for their impact. Barrier packages used packaging will vary with the nature of the product/package combination.

packaging in the Netherlands demonstrated that 'the packaging sub-system cannot be studied or optimised in isolation'. Accordingly, active packaging or processed foods at present are generally not autenable to economic recycling. The benefits of active packaging in terms of food quality, safety and shelf-life extension will need to be considered in a holistic approach to environmental impact assessment. A study by Kooijman (1994) of food plastics should be judged on the basis of their contribution to the quality and safety of food.

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Edible films and coatings as active layers S

B, CUQ, N. GONTARD and S. GUILBERT

5.1 Introduction

ipid films to protect meat products, and soy milk-based lipoprotein films to and conservation. The most common examples are wax coatings for fruit Edible films and coalings are traditionally used to improve food appearance (used in China since the 12th century), chocolate coatings for confectionery, improve the appearance and preservation of certain foods in Asia.

monotonous polymers, proteins have a specific structure which confers larger potential functional properties (Guilbert and Graille, 1994). Many ipidic compounds, such as animal and vegetable fats (natural waxes and derivalives, acetoplycerides, surface-active agents, etc.), have been used to make edible films and coatings (Guilbert and Biquet, 1989, Kester and Fennema, 1986). They are generally used for their excellent moisture barrier properties, but there can be problems concerning stability (particularly based films; this is due to the fact that, contrary to polysaccharides which are properties of these films are generally better than those of polysacchandecasein, etc. (Guilbert and Biquet, 1989). The mechanical and barrier proteins and lipidic compounds. Polysaccharides (vegetable and microbial gunts, starches, celluloses and derivatives, etc.) have good film-forming properties. Films formed from these hydrophilic compounds provide esticient barriers against oils and lipids (Munay et al., 1972), but their moisture barrier properties are poor. Although not as extensively studied. protein-based films have highly interesting properties. Many protein materials have been tested: collagen, zein, wheat gluten, ovalburain, soybean, Formulations for edible films or coatings must include at least one component able to form a suitably cohesive and continuous matrix. The basic materials can be classified in three categories: polysaccharidus,

overcome their respective drawbacks. Most composite films studied to date combine a lipidic compound and a hydrocolloid-based structural matrix (Cole, 1969; Darriels, 1973; Gornard et al., 1994a; Guilbert, 1986; Kamper Edible films and coatings formed with several compounds (composite functional properties of these different constitutive materials and to films) have been developed to take advantage of the complementary exidation), texture and organoleptic quality (opacity, waxy taste).

Coatings are formed directly on the food product using either liquid filmforming solutions (or dispersions) or molten compounds (e.g. lipids). They and Fennema, 1984a, b).

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